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U.S. APPLICE	3/647946	PCT/CA99/0029			1	94 MIS:jb	
21. The fol	lowing fees are submitted:.				CALCULATIONS	PTO USE ONLY	
BASIC NATIONA Neither interinter international	L FEE (37 CFR 1.492 (a) (1) - (mational preliminary examination search fee (37 CFR 1.445(a)(2) p lonal Search Report not prepared	fee (37 CFR 1.482) nor baid to USPTO	\$1,00	00.00			
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and an claim	ENTER APPROPRIA	\$1,000.00					
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Total claims	35 - 20 =	15	x \$18.0		\$270.00		
Independent claims	5 - 3 =	2	x \$78.0)()	\$156.00 \$0.00		
Multiple Dependen	t Claims (check if applicable). TOTAL OF	\$1,426.00					
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months from the earliest claimed priority date (37 CFR 1.492 (f)). TOTAL NATIONAL FEE = \$1,426.00 Fee for recording the enclosed assignment (37 CFR 1.21(h)). The assignment must be accompanied by an appropriate cover sheet (37 CFR 3.28, 3.31) (check if applicable). \$0.00							
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Mr. Michael I. St Sim & McBurney			SIGNAT	URE			
6th Floor, 330 Unviersity Avenue Toronto, Ontario Canada, M5G 1R7.			Michael I. Stewart NAME				
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PATENT TRADEMAK OFFICE				October 4, 2000			
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528 Rec'd PCT/PTO 0 6 OCT 2000

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Our Ref: 1038-1094 MIS:jb

In re National Phase of International Application

No.:

PCT/CA99/00292

International

Filing Date:

April 7, 1999

Applicant:

Robert C. Brunham

Title:

DNA IMMUNIZATION AGAINST CHLAMYDIA INFECTION

PRELIMINARY AMENDMENT

The Commissioner of Patents and Trademarks, Washington, D.C. 20231, U. S. A.

Dear Sir:

Please amend this application in the following manner:

In the Disclosure:

Before the first line of the specification, add the following:

" REFERENCE TO RELATED APPLICATIONS

This application is a national phase application under 35 U.S.C. 371 of PCT/CA99/00292."

REMARKS

The specification has been amended on page 1 to reflect that this application is a U.S. National Phase filing under 35 U.S.C. 371 of PCT/CA99/00292.

Respectfully submitted,

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Date: October 4, 2000

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TITLE OF INVENTION

DNA IMMUNIZATION AGAINST CHLAMYDIA INFECTION

FIELD OF INVENTION

The present invention relates to immunology and, in particular, to immunization of hosts using nucleic acid to provide protection against infection by Chlamydia.

BACKGROUND OF THE INVENTION

DNA immunization is an approach for generating protective immunity against infectious diseases (ref. 1 - throughout this application, various references are cited in parentheses to describe more fully the state of invention pertains. art to which this the bibliographic information for each citation is found at the end of the specification, immediately preceding the claims. The disclosure of these references are hereby incorporated by reference into the present disclosure). Unlike protein or peptide based subunit vaccines, DNA immunity provides protective immunization expression of foreign proteins by host cells, allowing the presentation of antigen to the immune system in a manner more analogous to that which occurs during infection with viruses or intracellular pathogens 2). Although considerable interest has generated by this technique, successful immunity has been most consistently induced by DNA immunization for Results have been more viral diseases (ref. 3). variable with non-viral pathogens which may reflect differences in the nature of the pathogens, immunizing antigens chosen, and in the routes Further development DNA immunization (ref. 4). vaccination will depend on elucidating the underlying immunological mechanisms and broadening its application for which other infectious diseases strategies of vaccine development have failed.

Chlamydia trachomatis is an obligate intracellular bacterial pathogen which usually remains localized to mucosal epithelial surfaces of the human host.

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Chlamydiae are dimorphic bacteria with an extracellular spore-like transmission cell termed the elementary body (EB) and an intracellular replicative cell termed the 5). From a public reticulate body (ref. are infections perspective, chlamydial importance because they are significant causes infertility, blindness and are a prevalent co-factor facilitating the transmission of human immunodeficiency Protective immunity to C. virus type 1 (ref. 6). trachomatis is effected through cytokines released by Th1-like CD 4 lymphocyte responses and by local antibody in mucosal secretions and is believed to be primarily directed to the major outer membrane protein (MOMP), which is quantitatively the dominant surface protein on the chlamydial bacterial cell and has a molecular mass of about 40 kDa (ref. 19).

Initial efforts in developing a chlamydial vaccine were based on parenteral immunization with the whole bacterial cell. Although this approach met with success in human trials, it was limited because protection was short-lived, partial and vaccination may exacerbate disease during subsequent infection episodes possibly due to pathological reactions to certain chlamydial antigens (ref. 8). More recent attempts at chlamydial vaccine design have been based on a subunit design using MOMP protein or peptides. These subunit vaccines have also generally failed, perhaps because the immunogens do not induce protective cellular and humoral immune responses recalled by native epitopes on the organism (ref. 9).

EP 192033 describes the provision of DNA construct for the expression, in vitro, of Chlamydia trachomatis MOMP polypeptides comprising the following operably linked elements:

35 a transcriptional promoter,

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a DNA molecule encoding a *C. trachomatis* MOMP polypeptide comprising a MOMP polynucleotide at least 27 base pairs in length from a sequence provided in Appendix A thereto, and

a transcriptional terminator, wherein at least one of the transcriptional regulatory elements is not derived from *Chlamydia trachomatis*. There is no disclosure or suggestion in this prior art to effect DNA immunization with any such constructs.

WO 94/26900 describes the provision of hybrid picornaviruses which express chlamydial epitopes from MOMP of *Chlamydia trachomatis* and which is capable of inducing antibodies immuno-reactive with at least three different *Chlamydia* serovars. The hybrid picornavirus preferably is a hybrid polio virus which is attenuated for human administration.

WO 98/02546, assigned to the assignee hereof and the disclosure of which is incorporated herein by reference, describes the DNA immunization of a host by a plasmid vector comprising a nucleotide sequence encoding a major outer membrane protein (MOMP) of a strain of Chlamydia or encoding the N-terminal half of MOMP.

SUMMARY OF THE INVENTION

The present invention is concerned with nucleic acid immunization, specifically DNA immunization, to generate in a host protective antibodies to a fragment of MOMP of a strain of *Chlamydia* that encompasses epitopic sequences. DNA immunization induces a broad spectrum of immune responses including Th1-like CD4 responses and mucosal immunity.

In one aspect of the invention, there is provided a non-replicating vector, comprising a nucleotide sequence encoding a region comprising at least one of the conserved domains 2, 3 and 5 of a major outer membrane protein of a strain of *Chlamydia*, and a promoter sequence operatively coupled to the nucleotide sequence

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for expression of the at least one conserved domain in a host.

A MOMP gene fragment that encompasses epitopic sequences may include one or more conserved domain (CD) sequences and/or one or more variable domain (VD) sequences of MOMP from a strain of Chlamydia. particular, the fragment may encompass the CD2 and VD2 sequences, CD3 and VD3 sequences and CD5 sequence. Clones containing nucleotide sequences encoding such fragments are termed clones CV2, CV3 and CD5 herein. Clone CV2 encompasses nucleotides 247 to Chlamydia trachomatis MOMP gene, clone CV3 encompasses nucleotides 469 to 696 of Chlamydia trachomatis MOMP gene and clone CV5 encompasses nucleotides 931 to 1098 gene. The Chlamydia trachomatis MOMP invention employs the conserved domains 2, 3 and 5.

The strain of *Chlamydia* may be a strain of *Chlamydia* inducing chlamydial infection of the lung, including *Chlamydia trachomatis* or *Chlamydia pneumoniae*. The non-replicating vector may be plasmid pcDNA3 into which the nucleotide sequence is inserted. The immune response which is stimulated may be predominantly a cellular immune response.

In one aspect of the present invention, there is provided an immunogenic composition for in vivo administration to a host for the generation in the host of a protective immune response to a major outer membrane protein (MOMP) of a strain of Chlamydia, comprising a non-replicating vector that generates a MOMP-specific immune response, and a promoter sequence operatively coupled to the nucleotide sequence for expression of the MOMP fragment in the host; and a pharmaceutically-acceptable carrier therefor.

In a further aspect of the invention, there is provided as a method of immunizing a host against disease caused by infection with a strain of *Chlamydia*,

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which comprises administering to the host an effective amount of a non-replicating vector as provided herein that generates a MOMP-specific immune response, and a promoter sequence operatively coupled to the nucleotide sequence for expression of the conserved sequence in the host.

In these aspects of the present invention, the various options and alternatives discussed above may be employed.

The non-replicating vector may be administrated to the host, including a human host, in any convenient manner, such as intramuscularly or intranasally. Intranasal administration stimulated the strongest immune response in experiments conducted herein.

includes, invention also in present a method of using aspect thereof, additional MOMP fragment sequence encoding a nucleotide generates a MOMP-specific immune response, to produce an immune response in a host, which comprises isolating the nucleotide sequence as described above, operatively linking the nucleotide sequence to at least one control sequence to produce a non-replicating vector, the control sequence directing expression of the MOMP fragment when introduced into a host to produce an immune response to the MOMP fragment, and introducing the vector into a host.

A further aspect of the present invention provides a method of producing a vaccine for protection of a host against disease caused by infection with a strain of Chlamydia, which comprises isolating a nucleotide sequence encoding a MOMP fragment as described above and that generates a MOMP-specific immune response, operatively linking the nucleotide sequence to at least one control sequence to produce a non-replicating vector, the control sequence directing expression of the MOMP fragment when introduced to a host to produce an

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immune response to the MOMP fragment, and formulating the vector as a vaccine for *in vivo* administration to a host. The invention extends to the vaccine produced by this method.

Advantages of the present invention, therefore, include a method of obtaining a protective immune response to infection carried by a strain of *Chlamydia* by nucleic acid immunization of nuelcic acid sequence encoding epitopic sequences of the major outer membrane protein of a strain of *Chlamydia* that generate a MOMP-specific immune response.

BRIEF DESCRIPTION OF DRAWINGS

Figure 1 shows the elements and construction of plasmid pcDNA3/MOMP, 6495 bp in size.

Figure 2 shows schematically the nucleotide structure of the mature MOMP gene of *C. trachomatis* MoPn strain with conserved (CD) and variable (VD) domains identified as well as clones formed by cloning the identified sequences into pcDNA3, as described below in the Examples.

Figure 3 shows the loss in body weight (in grams) following intranasal challenge with 5 x 10^3 IFU of MoPn among groups of Balb/c mice intramuscularly immunized with blank vector (pcDNA3), with pcDNA3 into which is individually cloned CV1 to CD5 encoding MOMP nucleotide sequences (CV1 etc), and with pcDNA3 into which the whole MOMP encoding nucleotide sequence is cloned (pMOMP):

Figure 4 shows the results of assays to determine growth of C. trachomatis on day 10 in lungs of mice challenged with 5 x 10^3 IFU of MoPn following intramuscular immunization with blank vector (pcDNA3), with pcDNA3 into which is individually cloned CV1 to CD5 encoding MOMP nucleotide sequences (pCV1 etc), and with pcDNA3 into which the whole MOMP encoding nucleotide sequence is cloned (pMOMP).

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Figure 5 shows footpad swelling reactions (DTH) 48 hours after footpad injection of 2×10^5 IFU of inactivated MoPn EBs among groups of Balb/c mice intramuscularly immunized with blank pcDNA3 vector (PC), with pcDNA3 into which is individually cloned CV1 to CD5 encoding MOMP nucleotide sequences (CV1 etc), and with pcDNA3 into which the whole MOMP encoding nucleotide sequence is cloned (pM).

Figure 6 shows the proliferation responses of splenocytes at day 60 post immunization after *in vitro* stimulation with whole inactivated MoPn EBs for 96 hours among groups of Balb/c mice immunized with blank pcDNA3 vector (pc), with pcDNA3 into which is individually cloned CV1 to CD5 encoding MOMP nucleotide sequences (CV1 etc), and with pcDNA3 into which the whole MOMP encoding nucleotide sequences is cloned (pM).

Figure 7 shows the proliferation responses of splenocytes to the same constructs is in Figure 6, except that the results are expressed as a stimulation index (SI).

Figure 8 shows the interferon- γ secretion response of MoPn stimulated splenocytes collected on day 60 after immunization among groups of Balb/c mice immunized with blank pcDNA3 vector (pc), with pcDNA3 into which is individually cloned CV1 to CD5 encoding MOMP nucleotide sequences (CV1 etc), and with pcDNA3 into which the whole MoPn MOMP encoding nucleotide sequence is cloned (pM).

Figure 9 shows the IgG2a antibody titer to whole MoPn EBs using sera collected at day 60 after immunization among groups of Balb/c mice immunized with blank pcDNA3 vector (pc), with pcDNA3 into which is individually cloned CV1 to CD5 encoding MOMP nucleotide sequences (CV1 etc), and with pcDNA3 into which the whole MOMP encoding nucleotide sequences is cloned (pM).

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Figure 10 shows a comparison of the amino acid sequence of MOMP sequences (SEQ ID NOS: 1 to 15) from a variety of serovars of *C. trachomatis*. Residues which are identical to serovar E MOMP are represented by dots. The four VDs (VDI to VDIV) and the conserved cysteines are boxed by solid line. The conserved position where one cysteine is located in all *C. trachomatis* and *C. pneumonitis* MOMP sequences, but where one serine is located in GPIC and Mn MOMPs, is boxed by a broken line. Numbers above boxes denote amino acid residues of serovar E MOMP only.

GENERAL DESCRIPTION OF THE INVENTION

To illustrate the present invention, plasmid DNA was constructed containing the MOMP gene fragments from the C. trachomatis mouse pneumonitis strain (MoPn), which is a natural murine pathogen, permitting experimentation to be effected in mice. It is known that primary infection in the model induces strong protective immunity to reinfection. For immunization, a human pathogen strain is used, such as serovar C of C. trachomatis.

Any convenient plasmid vector may be used for the MOMP gene fragment, such as pcDNA3, a eukaryotic IIselectable expression vector (Invitrogen, San Diego, CA, USA), containing a cyotmegalovirus promoter. The MOMP gene fragment may be inserted in the vector in any convenient manner. The gene fragments may be amplified from Chlamydia trachomatic genomic DNA by PCR using suitable primers and the PCR product cloned into the vector. The MOMP gene-carrying plasmid may transferred, such as by electroporation, into E. coli for replication therein. Plasmids may be extracted from the E. coli in any convenient manner.

The plasmid containing the MOMP gene fragment may be administered in any convenient manner to the host, such as intramuscularly or intranasally, in conjunction

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with a pharmaceutically-acceptable carrier. In the experimentation outlined below, it was found that intranasal administration of the plasmid DNA elicited the strongest immune response.

The data presented herein and described in detail below demonstrates that DNA immunization with specific C. trachomatis MOMP gene fragments elicits both cellular and humoral immune responses and produces significant protective immunity to lung challenge infection with C. trachomatis MoPn. The results are more encouraging than those obtained using recombinant MOMP protein or synthetic peptides as the immunogen and suggest that DNA immunization is an alternative method to deliver a chlamydial subunit immunogen in order to elicit the protective cellular requisite and humoral immune responses.

The data presented herein also demonstrate the importance of selection of an antigen gene fragment for DNA immunization. As described in the aforementioned WO 98/02546, the antigen gene elicits immune responses that are capable of stimulating recall immunity following In exposure to the natural pathogen. particular, injection of a DNA expression vector encoding the major surface protein (pMOMP) or fragment thereof but not one encoding a cytoplasmic enzyme (CTP synthetase) of C. trachomatis, generated significant protective immunity The protective subsequent chlamydial challenge. immune response appeared to be predominantly mediated by cellular immunity and not by humoral immunity since antibodies elicited by DNA vaccination did not bind to MOMP DNA but EBs. In addition, synthetase DNA immunization elicited cellular immunity readily recalled by native EBs as shown by positive DTH reactions.

In addition, as set forth in WO 98/02546, mucosal delivery of MOMP DNA is significantly more efficient in

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WO 99/51745 PCT/CA99/00292

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inducing protective immunity to C. trachomatis infection than intramuscular injection. This may be relevant to nature of *C. trachomatis* infection which essentially restricted to mucosal surfaces and the efficiency of antigen presentation (ref. 14). population and rapid recruitment of dendritic cells into the respiratory epithelium of the lung may be relevant to the enhanced efficacy of intranasal DNA immunization experiments (ref. 15). The data presented in WO 98/02546 demonstration of а first represents the subunit chlamydial vaccine which engenders substantial protective immunity.

Additionally, it may be possible to amplify (and/or canalize) the protective immune response by co-administration of DNAs that express immunoregulatory cytokines in addition to the antigen gene in order to achieve complete immunity (ref. 21) The use of multiple antigen genes from chlamydiae may augment the level of protective immunity achieved by DNA vaccination.

A possible concern regarding MOMP DNA immunization according to WO 98/02546 stems from the observation that the MOMP among human C. trachomatis strains is highly polymorphic (ref. 16) and hence it may be difficult to generate a universal chlamydial vaccine based on this One way to solve this problem is to antigen gene. search for conserved protective epitope(s) within the MOMP molecule, as described herein. As seen in the results presented below, certain vectors containing nucleotide sequences encoding conserved and variable domains, identified in Figure 2, or conserved domains generated a protective immune response, as determined by loss of body weight, as shown in Figure 3. Figure 4 shows that the pCV3 and pCD5 immunogen evoked a protective immune response to MoPn challenge as measured by in vivo growth of MoPn in lung tissue day 10 post challenge and comparable to pMOMP. Figure 5 shows that

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immunization with the vectors elicited variable positive DTH responses for footpad injection of MoPn Ebs.

Figures 6 and 7 show the proliferation responses of splenocytes to the vectors containing the conserved and variable domains and the whole MOMP gene. The results set forth in Figures 6 and 7 show that pCV3 and pMOMP elicit a cell mediated immune response.

Figure 8 shows interferon- γ secretion responses of the splenocytes to the vectors containing the conserved and variable domains and the whole MOMP gene. The results obtained in Figure 8 suggest that cytokine generation may not necessarily be a correlate of a protective immune response.

Another, possibly more feasible, way is to design a multivalent vaccine based on multiple MOMP genes. The latter approach is justified by the fact that the inferred amino acid sequences of MOMP among related serovars is relatively conserved (see Figure 10) and the repertoire of *C. trachomatis* gene variants appears to be finite (ref. 16).

It is clearly apparent to one skilled in the art, that the various embodiments of the present invention have many applications in the fields of vaccination, diagnosis and treatment of chlamydial infections. A further non-limiting discussion of such uses is further presented below.

1. Vaccine Preparation and Use

Immunogenic compositions, suitable to be used as vaccines, may be prepared from the MOMP gene fragments thereof and vectors as disclosed herein. The vaccine elicits an immune response in a subject which includes the production of anti-MOMP antibodies. Immunogenic compositions, including vaccines, containing the nucleic acid may be prepared as injectables, in physiologically-acceptable liquid solutions or emulsions for polynucleotide administration. The nucleic acid may be

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associated with liposomes, such as lecithin liposomes or other liposomes known in the art, as a nucleic acid liposome (for example, as described in WO 93/24640) or the nucleic acid may be associated with an adjuvant, as described in more detail below. Liposomes comprising cationic lipids interact spontaneously and rapidly with polyanions, such as DNA and RNA, resulting liposome/nucleic acid complexes that capture up to 100% of the polynucleotide. In addition, the polycationic complexes fuse with cell membranes, resulting in an intracellular delivery of polynucleotide that bypasses the degradative enzymes of the lysosomal compartment. WO 94/27435 describes compositions for genetic immunization comprising cationic lipids and polynucleotides. which assist in the cellular uptake of nucleic acid, calcium ions, viral proteins such as and transfection facilitating agents, may advantageously be used.

Polynucleotide immunogenic preparations may also be formulated as microcapsules, including biodegradable time-release particles. Thus, U.S. Patent 5,151,264 describes a particulate carrier of phospholipid/glycolipid/polysaccharide nature that has been termed Bio Vecteurs Supra Moléculaires (BVSM). The particulate carriers are intended to transport a variety of molecules having biological activity in one of the layers thereof.

U.S. Patent 5,075,109 describes encapsulation of the antigens trinitrophenylated keyhole limpet hemocyanin and staphylococcal enterotoxin B in 50:50 (DL-lactideco-glycolide). Other polymers encapsulation are suggested, such as poly(glycolide), poly(DL-lactide-coglycolide), copolyoxalates, poly(lactide-co-caprolactone), polycaprolactone, poly(esteramides), polyorthoesters and poly(8hydroxybutyric acid), and polyanhydrides.

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WO 91/06282 describes a delivery vehicle comprising a plurality of bioadhesive microspheres and antigens. The microspheres being of starch, gelatin, dextran, collagen or albumin. This delivery vehicle is particularly intended for the uptake of vaccine across the nasal mucosa. The delivery vehicle may additionally contain an absorption enhancer.

The MOMP gene fragment containing non-replicating vectors may be mixed with pharmaceutically acceptable excipients which are compatible therewith. excipients may include, water, saline, and combinations thereof. glycerol, ethanol, immunogenic compositions and vaccines further may contain auxiliary substances, such as wetting or emulsifying agents, pH buffering agents, or adjuvants to the effectiveness thereof. Immunogenic enhance administered compositions and vaccines may be injection subcutaneously, by parenterally, intradermally or intramuscularly, intravenously, possibly following pretreatment of the injection site with a local anesthetic. Alternatively, the immunogenic compositions formed according to the present invention, may be formulated and delivered in a manner to evoke an response at mucosal surfaces. Thus, immune immunogenic composition may be administered to mucosal surfaces by, for example, the nasal or (intragastric) routes. Alternatively, other modes of administration including suppositories and formulations may be desirable. For suppositories, and carriers include, for example, binders may triglycerides. polyalkylene glycols or formulations may include normally employed incipients, example, pharmaceutical grades of as, for saccharine, cellulose and magnesium carbonate.

The immunogenic preparations and vaccines are administered in a manner compatible with the dosage

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in such amount will be formulation, and as therapeutically effective, protective and immunogenic. The quantity to be administered depends on the subject to be treated, including, for example, the capacity of the individual's immune system to synthesize the MOMP and antibodies thereto, and if needed, to produce a cell-mediated immune response. Precise amounts active ingredient required to be administered depend on the judgement of the practitioner. However, suitable dosage ranges are readily determinable by one skilled in the art and may be of the order of about 1 μg to about 1 mg of the MOMP gene-containing vectors. Suitable regimes for initial administration and booster doses are also variable, but may include an initial administration followed by subsequent administrations. The dosage may also depend on the route of administration and will vary according to the size of the host. A vaccine which protects against only one pathogen is a monovalent vaccine. Vaccines which contain antigenic material of several pathogens are combined vaccines and also belong the present invention. Such combined vaccines contain, for example, material from various pathogens or from various strains of the same pathogen, or from combinations of various pathogens.

Immunogenicity can be significantly improved if the vectors are co-administered with adjuvants, commonly used as 0.05 to 0.1 percent solution in phosphate-buffered saline. Adjuvants enhance the immunogenicity of an antigen but are not necessarily immunogenic themselves. Adjuvants may act by retaining the antigen locally near the site of administration to produce a depot effect facilitating a slow, sustained release of antigen to cells of the immune system. Adjuvants can also attract cells of the immune system to an antigen depot and stimulate such cells to elicit immune responses.

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Immunostimulatory agents or adjuvants have been used for many years to improve the host immune responses to, for example, vaccines. Thus, adjuvants have been identified that enhance the immune response to antigens. Some of these adjuvants are toxic, however, and can cause undesirable side-effects, making them unsuitable for use in humans and many animals. Indeed, only aluminum hydroxide and aluminum phosphate (collectively commonly referred to as alum) are routinely used as adjuvants in human and veterinary vaccines.

A wide range of extrinsic adjuvants and other immunomodulating material can provoke potent immune responses to antigens. These include saponins complexed membrane protein antigens to produce stimulating complexes (ISCOMS), pluronic polymers with killed mycobacteria in mineral mineral oil, Freund's complete adjuvant, bacterial products, such as muramyl dipeptide (MDP) and lipopolysaccharide (LPS), as well as Quil A derivatives and components thereof, QS hydroxide, zinc phosphate, calcium calcium an octodecyl ester of an amino acid, hydroxide, polyphosphazene. ISCOPREP, DC-chol, DDBA and Advantageous combinations of adjuvants are described in Applications Patent United States copending 08/261,194 filed June 16, 1994 and 08/483,856 filed June 7, 1995, assigned to the assignee hereof and the disclosures of which are incorporated herein by reference thereto (WO 95/34308).

In particular embodiments of the present invention, the non-replicating vector comprising a first nucleotide sequence encoding a MOMP gene fragment of *Chlamydia* may be delivered in conjunction with a targeting molecule to target the vector to selected cells including cells of the immune system.

The non-replicating vector may be delivered to the host by a variety of procedures, for example, Tang et

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al. (ref. 17) disclosed that introduction of gold microprojectiles coated with DNA encoding bovine growth hormone (BGH) into the skin of mice resulted in production of anti-BGH antibodies in the mice, while Furth et al. (ref. 18) showed that a jet injector could be used to transfect skin, muscle, fat and mammary tissues of living animals.

2. Immunoassays

The MOMP gene fragments and vectors of the present invention also are useful as immunogens for antibodies in anti-MOMP for use generation of immunoassays, including enzyme-linked immunosorbent RIAs and other non-enzyme assays (ELISA), antibody binding assays or procedures known in the art. In ELISA assays, the non-replicating vector first is administered to a host to generate antibodies specific These MOMP specific antibodies are to the MOMP. immobilized onto a selected surface, for example, a surface capable of binding the antibodies, such as the wells of a polystyrene microtiter plate. After washing incompletely adsorbed antibodies, remove nonspecific protein, such as a solution of bovine serum albumin (BSA) that is known to be antigenically neutral with regard to the test sample, may be bound to the for selected surface. This allows blocking nonspecific adsorption sites on the immobilizing surface and thus reduces the background caused by nonspecific bindings of antisera onto the surface.

The immobilizing surface is then contacted with a sample, such as clinical or biological materials, to be tested in a manner conducive to immune complex (antigen/antibody) formation. This procedure may include diluting the sample with diluents, such as solutions of BSA, bovine gamma globulin (BGG) and/or phosphate buffered saline (PBS)/Tween. The sample is then allowed to incubate for from about 2 to 4 hours, at

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temperatures such as of the order of about 20° to 37°C. Following incubation, the sample-contacted surface is washed to remove non-immunocomplexed material. The washing procedure may include washing with a solution, such as PBS/Tween or a borate buffer. Following formation of specific immunocomplexes between the test sample and the bound MOMP specific antibodies, and subsequent washing, the occurrence, and even amount, of immunocomplex formation may be determined.

10 EXAMPLES

above disclosure generally describes present invention. A more complete understanding can be reference to the following by obtained These Examples are described solely for purposes of illustration and are not intended to limit the scope of the invention. Changes in form and contemplated substitution of equivalents are circumstances may suggest or render expedient. Although specific terms have been employed herein, such terms are intended in a descriptive sense and not for purposes of limitation.

Example 1:

This Example illustrates the preparation of a plasmid vector containing the MOMP gene, as also described in WO 98/02546.

pMOMP expression vector was made as follows. MOMP gene was amplified from Chlamydia trachomatis mouse pneumonitis (MoPn) strain genomic DNA by polymerase 5′ primer reaction (PCR) with a chain (GGGGATCCGCCACCATGCTGCCTGTGGGGAATCCT) (SEQ ID NO: which includes a BamHl site, a ribosomal binding site, an initiation codon and the N-terminal sequence of the 3' mature MOMP of MoPn and (GGGGCTCGAGCTATTAACGGAACTGAGC) (SEQ ID NO: 17) which includes the C-terminal sequence of the MoPn MOMP, a Xhol site and a stop codon. The DNA sequence of the MOMP

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leader peptide gene sequence was excluded. After digestion with BamH1 and Xhol, the PCR product was II-selectable eukaryotic into the pcDNA3 Diego) vector (Invitrogen, San expression transcription under control of the human cytomegatovirus major intermediate early enhancer region (CMV promoter). The MOMP gene-encoding plasmid was transferred by electroporation into $E.\ coli$ DH5lphaF which was grown in LB broth containing 100 μ g/ml of ampicillin. The plasmids was extracted by $Wizard^{TM}$ Plus Maxiprep DNA purification sequence of The system (Promega, Madison). MOMP gene was verified by PCR direct recombinant sequence analysis, as described (ref. 20). plasmid DNA was dissolved in saline at a concentration of 1 mg/ml. The DNA concentration was determined by a DU-62 spectrophotometer (Beckman, Fullerton, CA) at 260 nm and the size of the plasmid was compared with DNA standards in ethidium bromide-stained agarose gel.

The MOMP gene containing so obtained plasmid, pcDNA3/MOMP, and its constitutive elements are shown in Figure 1. A similar plasmid (pM(C)) was constructed from the MOMP gene serovar C of $C.\ trachomatis$.

For experimental design, groups of 4 to 5 week old female Balb/c mice (5 to 13 per group) were immunized intramuscularly (IM) or intranasally (IN) with plasmid DNA containing the coding sequence of the MoPn MOMP gene (1095 bp), prepared as described in Example 1, or with the coding sequence of the C. trachomatis serovar L_2 CTP synthetase gene (1619 bp (refs. 10, 12), prepared by a procedure analogous described in Example 1. CTP synthetase is a conserved chlamydial cytoplasmic enzyme catalizing the final step in pyrimidine biosynthesis and is not known to induce protective immunity. Negative control animals were injected with saline or with the plasmid vector lacking an inserted chlamydial gene.

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Example 2:

This Example illustrates DNA immunization of mice and the results of DTH testing.

A model of murine pneumonia induced by the *C. trachomatis* mouse pneumonitis strain (MoPn) was used (ref. 11). Unlike most strains of *C. trachomatis* which are restricted to producing infection and disease in humans, MoPn is a natural murine pathogen. It has previously been demonstrated that primary infection in this model induces strong protective immunity to reinfection. In addition, clearance of infection is related to CD4 Th1 lymphocyte responses and is dependent on MHC class II antigen presentation (ref. 11).

For IM immunization, both quardiceps were injected with 100 μ g DNA in 100 μ l of saline per injection site on three occasions at 0, 3 and 6 weeks. For IN immunization, anaesthetized mice aspirated 25 μ l of saline containing 50 μ g DNA on three occasions at 0, 3 and 6 weeks. As a positive control, a separate group of mice received 5 x 10⁶ inclusion forming units (IFUs) of MoPn EBs administered intraperitoneally in incomplete Freund's adjuvant according to the above schedule. At week 8, all groups of mice had sera collected for measuring antibodies and were tested for delayed-type hypersensitivity (DTH) to MoPn Ebs by footpad injection (ref. 13).

A positive 48 and 72 hour DTH reaction was detected among mice immunized with MOMP DNA or with MoPn Ebs but not among mice immunized with the blank vector (see Figure 1 of WO 98/02546). The DTH reaction elicited with MOMP DNA delivered intranasally was comparable to that observed among mice immunized with EBs. No DTH reaction was detected among the groups of mice vaccinated with CTP synthetase DNA (see Table 1 below). Thus, injection of MOMP DNA generated a DTH reaction that was capable of

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recall by naturally processed peptides from C. trachomatis EBs while injection of CTP synthetase DNA failed to do so.

Example 3:

This Example illustrates DNA immunization of mice and the generation of antibodies.

Injection of CTP synthetase DNA as described in Example 2 resulted in the production of serum antibodies to recombinant CTP synthetase (Table 1) (ref. Antigen-specific serum Abs were measured by ELISA. Flatbottom 96-well plates (Corning 25805, Corning Science Products, Corning, NY) were coated with recombinant chlamydial CTP-synthetase (1 μg/ml) purified MoPn EBs (6 x 10^4 IFU/well) overnight at 4° C. The Plates were rinsed with distilled water and blocked with 4% BSA PBS-Tween and 1% low fat skim milk for 2 hours at room temperature. Dilutions of sera samples performed in 96-well round bottom immediately prior to application on the antigen coated plates. The plates were incubated overnight at 4°C and washed ten times. Biotinylated goat anti-mouse IgG1 or Biotechnology anti-mouse IgG2a (Southern Associates, Inc. Birmingham, AL) were next applied for 1 hour at 37°C. After washing, streptoavidin-alkaline ImmunoResearch phosphatase (Jackson conjugate Laboratories, Inc. Mississagua, Ontario, Canada) were added and incubated at 37°C for 30 min. Following another wash step, phosphatase substrate in phosphatase buffer (pH 9.8) was added and allowed to develop for 1 hour. The plates were read at 405 nm on a BIORAD 3550 microplate reader.

IgG2a antibody titers were approximately 10-fold higher than lgG1 antibody titers suggesting that DNA immunization elicited a more dominant $T_{\rm H1}$ -like response. Injection of MOMP DNA as described in Example 2 resulted

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in the production of serum antibodies to MOMP (Table 2) as detected in an immunoblot assay (Figure 2 of WO 98/02546). However, neither CTP synthetase DNA nor MOMP DNA immunized mice produced antibodies that bound to native *C. trachomatis* EBs (Table 1), suggesting that the antibody responses may not to be the dominantly protective mechanism.

Example 4:

This Example illustrates DNA immunization of mice to achieve protection.

a cell-mediated To investigate whether response elicited by MOMP DNA was functionally significant, in vivo protective efficacy was evaluated in mice challenged intranasally with 1 x 10^3 IFU of C. trachomatis MoPn. To provide a measure of Chlamydiainduced morbidity, the loss in body weight was measured over 10 days following challenge with C. trachomatis. Mice injected with the unmodified vector were used as negative controls and mice immunized with EBs were used as positive controls. Mice immunized with MOMP intranasally maintained a body weight comparable to that observed among EB immunized mice. Mice intramuscularly immunized with MOMP DNA lost body mass but did so at a rate less than the negative control group.

A more direct measure of the effectiveness of DNA vaccination is the ability of mice immunized with MOMP DNA to limit the *in vivo* growth of Chlamydia following a sublethal lung infection. Day 10 post-challenge is the time of peak growth (ref. 13) and was chosen for comparison of lung titers among the various groups of mice. Mice intranasally immunized with MOMP DNA had chlamydial lung titers that were over 1000-fold lower (\log_{10} IFU 1.3 ± 0.3 ; mean \pm SEM) than those of control mice immunized with the blank vector (\log_{10} IFU 5.0 ± 0.3 ; p<0.01). Mice intramuscularly immunized with MOMP DNA had chlamydial lung titers that were more than 10-fold

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lower than the unmodified vector group (p = 0.01). Mice intranasally immunized with MOMP DNA had significantly lower chlamydial lung titers than mice immunized with MOMP DNA intramuscularly (log10 IFU 1.3±0.8 versus log10 IFU 0.66 ± 0.3 respectively; p = 0.38). The substantial difference (2.4 logs) in chlamydial lung titers observed between the intranasally and intramuscularly MOMP DNA immunized mice suggests that mucosal immunization is efficient at inducing immune responses accelerate chlamydial clearance in the lung. The lack of protective effect with the unmodified vector control confirms that DNA per se was not responsible for the immune response. Moreover, the absence of protective immunity following immunization with CTP synthetase DNA confirms that the immunity was specific to the MOMP DNA (see Table 1).

Example 5:

This Example describes the construction of plasmids containing fragments of MOMP DNA.

A series of vectors was generated following the procedure outlined in Example 1 containing fragments of the nucleotide sequence of the MoPn MOMP gene by PCR cloning and subsequent cloning into the vector pcDNA3 to generate plasmids pCV1, pCV2, pCV3, pCV4 and pCD5, respectively, containing the respective fragments of the MoPn MOMP gene shown in Figure 2.

Example 5:

This Example illustrates immunization of mice with pCV1, pCV2, pCV3, pCV4 and pCD5.

Balb/c mice were immunized in the quadriceps three times at a three week intervals with 100 μg of pCV1, pCV2, pCV3, pCV4 and pCD5 DNA, following the procedure described in Example 2.

Fifteen days after the last immunization and 60 days after the first injection, mice were bled for

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measurement of serum antibodies of MoPn EBs in an EIA assay and were injected in the footpad with 25 μ l (5 x 10⁴ inclusion forming units) of heat killed EBs for measurement of DTH which was measured at 72 hours (ref. 13). Mice were intranasally challenged with 1000 infectious units of MoPn and their body weight measured daily for the subsequent 10 days. At that time, mice were sacrificed and quantitative cultures of MoPn in the lung determined (ref. 13).

Figure 3 shows that pCV2, pCV3 and pCD5 immunization evoked a protective immune response to MoPn challenge as measured by loss in body weight post infection comparable to that in mice protected against disease. Figure 4 shows that pCV3 and pCD5 immunization evoked a protective immune response to MoPn challenge as measured by *in vivo* growth of MoPn in lung tissue, comparable to pMOMP.

However, the specific domains eliciting these immune responses do not include those predicted in the art to contain T-cell epitopes. In this regard, several groups have attempted to define MOMP T-cell epitopes (refs. 22 to 26). All of those studies used overlapping synthetic peptides to various regions of the MOMP protein to prime mice. None of the predicted epitopes fall within regions that have been found to be protective.

Figure 5 shows that immunization with pCV1, pCV2, pCV3, pCV4 and pCD5 elicited variable positive DTH responses to footpad injection of MoPn EBs. pCV3 and pCD5 elicited greater responses, comparable to pMOMP. Immunization with the unmodified vector elicited neither serum antibodies nor a DTH response.

Figure 9 shows IgG_{2a} antibody titers in sera collected from the mice 60 days post immunization by the vectors containing the conserved and variable domains and full length MOMP gene. Only in the case of

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immunization by pCV3 and pCD5, was an IgG_{2a} immune response generated, indicating that a Th1-like response was elicited by these vectors.

As may be seen in this Example, the vectors containing specific segments of the MOMP gene were able to protect against disease, based on body weight loss, namely pCV2 and pCD5. In addition, vectors pCV3 and pCD5 were able to protect against infection, based on lung titres.

10 Example 6

This Example illustrates the proliferation response of splenocytes to the vectors pMOMP, pCV1, pCV2, pCV3, pCV4 and pCD5.

Mice were sacrificed two weeks after the fourth immunization following the protocol of Example 2. The spleens were removed and single-cell suspensions were prepared. 200 μ l of the cell suspension (5 x 10⁵ well) in RPMI-1640 medium containing 10% heat-inactivated fetal calf serum (FCS), 1% L-glutamine and 5 x 10^{-5} M 2mercaptoethanol (2ME, Kodak, Rochester, NY) incubated with 1 x 10^5 IFU/ml of MoPn in 96 well flat bottom plates in triplicate 37°C in 5% CO2 for 96 hours. Negative control wells contained spleen cells without antigen and positive control wells contained spleen cells with 0.25 μ g/ml of concanavalin A. 0.25 μ Ci/well of tritiated (3H) thymidine (2 Ci/mmol, 74 Gbq/mmol, imCi/ml, ICN, Irvine, CA) was added after 3 days of culture and 16h before harvest. The cells harvested with PHD cell harvester (Cambridge а Technology Inc., Watertown, MA, USA) and counted in 2ml of scintillation solution (Universal, ICN, Costa Mesa) in a Beckman LS5000 counter (Beckman Instrument, UK).

As may be seen in the results presented into Figures 6 and 7, pCV3 and pMOMP elicited a cell-mediated immune response.

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Example 7

This Example illustrates the interferon- γ secretion responses of splenocytes to the vectors pMOMP, pCV1, pCV2, pCV3, pCV4 and pCD5.

A cytokine-specific ELISPOT assay was used for the quantification of murine IFN γ and IL-10 secreting cells in the murine spleen. For all assays 96-well nitrocellulose-based microtiters (Milititer Multiscreen HA plates, Millipore Corp, Molshem, France) were coated overnight at 4°C with 100 μ l of the anti-cytokine mAb diluted in PBS at a concentration of 5 μ g/ml. After removing the coating solution from the plates, wells were blocked for at least 1 hour with RPMI-1640 media containing 40% fetal calf serum at 37°C, in CO2-. After rinsing the plates with PBS-T once, the testing cells were added into the wells.

For induction of antigen specific IFNy secreting cells in immunized mice, single cells were adjusted to 5 \times 10⁶ cells/ml and cultured with 2 \times 10⁵ IFU/ml of UVkilled EB of MoPn in 24 well plates for 72 hours. After washing with RPMI 1640, cells were added onto the 96well plates for 72 hours. After washing with RPMI 1640, cells were added onto the 96-well nitrocellulose-based microtiter plates which had been previously coated with anti-cytokine antibodies. The cells were added individual wells (2 x 10^5 or 1 x $10^5/100-\mu$ l/well) and incubated for 24 hours at 37°C in a CO2 incubator. Wells were rinsed extensively with PBS-T containing 1% BSA. Following rinsing with PBS-T three times (removing the supporting manifold and washing the back of the plate thoroughly with PBS-T), alkaline phosphatase conjugated streptavidin in PBS containing 1% BSA at 1:2000 at a concentration of 0.5 μ g/ml was added and incubated at 37°C in CO₂ for 45 min. After rinsing thoroughly, 100

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 μ l/well of the colormetric substrate phosphate BICP (5-bromo-4-chloro-3-indolyl phosphate)/NBT (Nitro blue tetrazolium) at 0.16 mg/ml BICP and 1 mg/ml NBT in substrate buffer (0.1 M NaCl, 0.1M Tris, pH 9.5, 0.05 M MgCl₂) was added and incubated at room temperature until spots were visualized. The reaction was stopped by the addition of water.

The results obtained are set forth in Figure 8 and suggest that cytokine generation may not necessarily be a correlate of a protective immune response.

SUMMARY OF DISCLOSURE

In summary of this disclosure, the present invention provides a method of nucleic acid, including DNA, immunization of a host, including humans, against disease caused by infection by a strain of *Chlamydia*, specifically *C. trachomatis*, employing a non-replicating vector, specifically a plasmid vector, containing a nucleotide sequence encoding an epitopic fragment of a major outer membrane protein (MOMP) of a strain of *Chlamydia* which generates a MOMP-specific immune response, and a promoter to effect expression of the MOMP fragment in the host. Modifications are possible within the scope of this invention.

Table 1

Serum antibody titers and delayed-type hypersensitivity (DTH) responses and in vivo growth Results are of Chlamydia trachomatis following pCTP synthetase or MoPn EB immunization. presented as means ± SEM.

	Anti-MoPn EB antibodies (log10)	Pn EB s (log ₁₀)	anti-rCTP antibodie	anti-rCTP synthetase antibodies (log10)	Anti-EB DTH $(mm \times 10^2)$	Anti-EB DTH log_{10} IFU/lung (mm x 10^2) d10 post
	IgG1	IgG2a	IgG1	IgG2a		Cilatrelige
Saline $(n = 9)$	2	<2	2	<2	4.5 ± 1.5	4.9 ± 2.4
pCTP synthetase	<2	<2	3.8 ± .3	4.7 ± .1	1.4 ± 1.5	$4.7 \pm .13$
(n = 11)						
EB (n = 4)	5.0 ± .3	4.8 ± .3	1.3 4.8 ± .3 3.6 ± .8	2.9 ± 0	15.2 ± 2.0	0

Table 2

Serum antibody Elisa titers to *Chlamydia trachomatis* mouse pneumonitis recombinant MOMP and EBs were measured 60 days after the initial immunization among mice immunized with blank vector alone (pcDNA3), vector containing the MOMP gene (pMOMP) and vector containing the CTP synthetase gene (pCTP). Non-immunized mice were also tested.

	rN	IOMP	EB		
Immunogen	IgG2a	IgG1	IgG2a	IgG1	
pcDNA3	<2.6*	<2.6	<2.6	<2.6	
pMOMP	3.77 ± 0.1	2.90 ± 0.14	3.35 ± 0.11	<2.6	
pCTP	ND	ND	<2.6	<2.6	
Preimmunization	<2.6	<2.6	<2.6	<2.6	

^{*} \log_{10} mean \pm SE IgG isotype speciic antibody titer ND = not done

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What I claim is:

1. A non-replicating vector, comprising:

a nucleotide sequence encoding a region which is at least one of the conserved domains 2, 3 and 5 of a major outer membrane protein of a strain of Chlamydia, and

- a promoter sequence operatively coupled to said nucleotide sequence for expression of said at least one conserved domain in a host.
- 2. The vector of claim 1 wherein said nucleotide sequence encoding the conserved domain 2 and/or 3 further includes a nucleotide sequence encoding a variable domain of the major outer membrane protein immediately downstream of the conserved domain.
- 3. The vector of claim 1 wherein said nucleotide sequence encodes the conserved domain 5 of the outer membrane protein.
- 4. The vector of claim 1 wherein said promoter sequence is the cytomegalovirus promoter.
- 5. The vector of claim 1 wherein said non-replicating vector comprises plasmid pcDNA3 containing said promoter sequence and into wherein said nucleotide sequence is inserted in operative position to said promoter sequence.
- 6. The vector of claim 5 wherein said strain of Chlamydia is a strain producing chlamydial infectious of the lung.
- 7. The vector of claim 5 wherein said strain of Chlamydia is a strain of Chlamydia trachomatis.
- 8. An immunogenic composition for in vivo administration to a host for the generation in the host of a protective immune response to a fragment of a major outer membrane protein (MOMP) of a strain of Chlamydia, comprising a non-replicating vector comprising:

a nucleotide sequence encoding a region which is at least one of the conserved domains 2, 3 and 5 of a major

outer membrane protein of a strain of Chlamydia and that generates a MOMP-specific immune response, and

- a promoter sequence operatively coupled to said nucleotide sequence for expression of said MOMP or MOMP fragment in the host; and
 - a pharmaceurically-acceptable carrier therefor.
- 9. The immunogenic composition of claim 8 wherein said nucleotide sequence encoding the conserved domain 2 and/or 3 further includes a nucleotide sequence encoding a variable domain of the major outer membrane protein immediately downstream of said conserved domain.
- 10. The immunogenic composition of claim 8 wherein said nucleotide sequence encodes the conserved domain 5 of a major outer membrane protein of a strain of Chlamydia.
- 11. The immunogenic composition of claim 8 wherein said promoter sequence is the cytomegalovirus promoter.
- 12. The immunogenic composition of claim 1 wherein said strain of Chlamydia is a strain producing chlamydial infections of the lung.
- 13. The immunogenic of claim 8 wherein said strain of Chlamydia is a strain of Chlamydia trachomatis.
- 14. The immunogenic composition of claim 13 wherein said non-replicating vector comprises plasmid pcDNA3 containing said promoter sequence and into which said nucleotide sequence is inserted in operative relation to said promoter sequence.
- 15. The composition of claim 8 wherein said immune response is predominantly a cellular immune response.
- 16. A method of immunizing a host against disease caused by infection with a strain of Chlamydia, which comprises administering to said host an effective amount of a non-replicating vector comprising:
- a nucleotide sequence encoding a region which is at least one of the conserved domains 2, 3 and 5 of a major

outer membrane protein of a strain of Chlamydia and that generates a MOMP-specific immune response, and

- a promoter sequence operatively coupled to said nucleotide sequence for expression of said MOMP in the host.
- 17. The method of claim 16 wherein said promoter sequence is the cytomegalovirus promoter.
- 18. The method of claim 16 wherein said strain of chlamydia is a strain producing chlamydial infections of the lung.
- 19. The method of claim 16 wherein said strain of Chlamydia is a strain of Chlamydia trachomatis.
- 20. The method of claim 16 wherein said non-replicating vector comprises plasmid pcDNA3 containing said promoter into which said nucleotide sequence is inserted in operative relation to said promoter sequence.
- 21. The method of claim 16 wherein said immune response is predominantly a cellular immune response.
- 22. The method of claim 16 wherein said non-replicating vector is administered intranasally.
- 23. The method of claim 16 wherein said host is a human host.
- 24. A method of using a nucleotide sequence encoding a fragment of a major outer membrane protein (MOMP) of a strain of Chlamydia that generates a MOMP-specific immune response, to produce an immune response in a host, which comprises:

isolating said nucleotide sequence encoding a region which is at least one of the conserved domains 2, 3 and 5 of a major outer membrane protein of a strain of Chlamydia.

operatively linking said nucleotide sequence to at least one control sequence to produce a non-replicating vector, said control sequence directing expression of said

introducing said vector into a host.

- 25. The method of claim 24 wherein said nucleotide sequence encoding the conserved domain 2 and/or 3 further includes a nucleotide sequence encoding a variable domain of the major outer membrane protein immediately downstream of said conserved domain.
- 26. The method of claim 24 wherein said nucleotide sequence encodes the conserved domain 5 of a major outer membrane protein of a strain of Chlamydia.
- 27. The method of claim 24 wherein said control sequence is the cytomegalovirus promoter.
- 28. The method of claim 24 wherein said strain of Chlamydia is a strain producing chlamydial infections of the lung.
- 29. The method of claim 24 wherein said strain of Chlamydia is a strain of Chlamydia trachomatis.
- 30. The method of claim 24 wherein said non-replicating vector comprises plasmid pcDNA3 containing said control sequence into which said gene encoding MOMP is inserted in operative relation to said control sequence.
- 31. The method of claim 24 wherein said immune response is predominantly a cellular immune response.
- 32. The method of claim 24 wherein said vector is introduced into said host intranasally.
- 33. The method of claim 24 wherein said host is a human host.
- 34. A method of producing a vaccine for protection of a host against disease caused by infection with a strain of Chlamydia, which comprises:

isolating a nucleotide sequence encoding a a region which is at least one of the conserved domains 2, 3 and 5 of a major outer membrane protein of a strain of Chlamydia and that generates a MOMP-specific immune response,

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operatively linking said nucleotide sequence to at least one control sequence to produce a non-replicating vector, the control sequence directing expression of said MOMP fragment when introduced to a host to produce an immune response to said MOMP fragment, and

formulating said vector as a vaccine for in vivo administration to a host.

35. A vaccine produced by the method of claim 34.

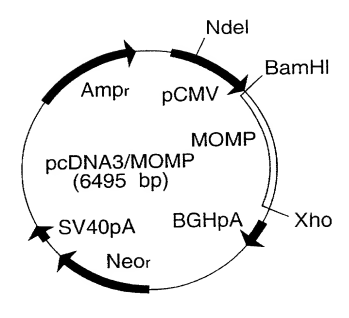
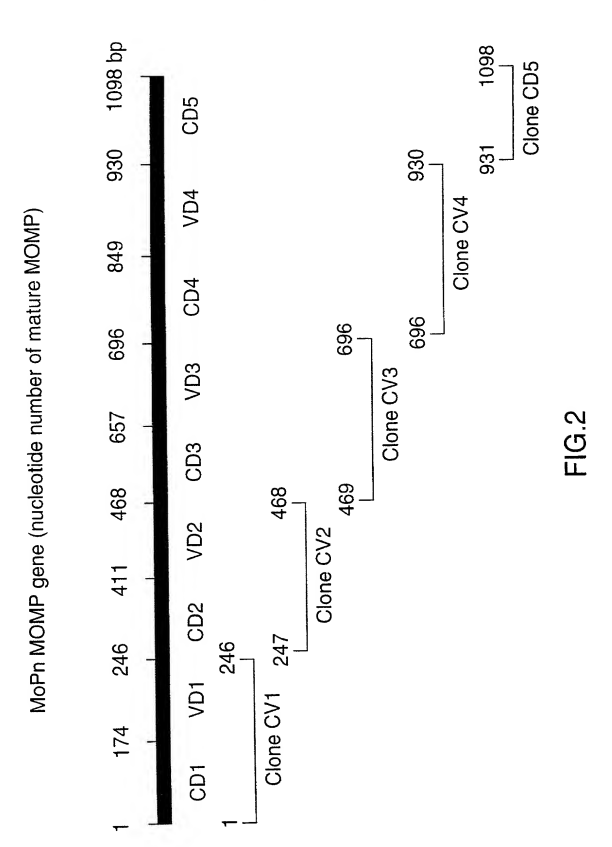
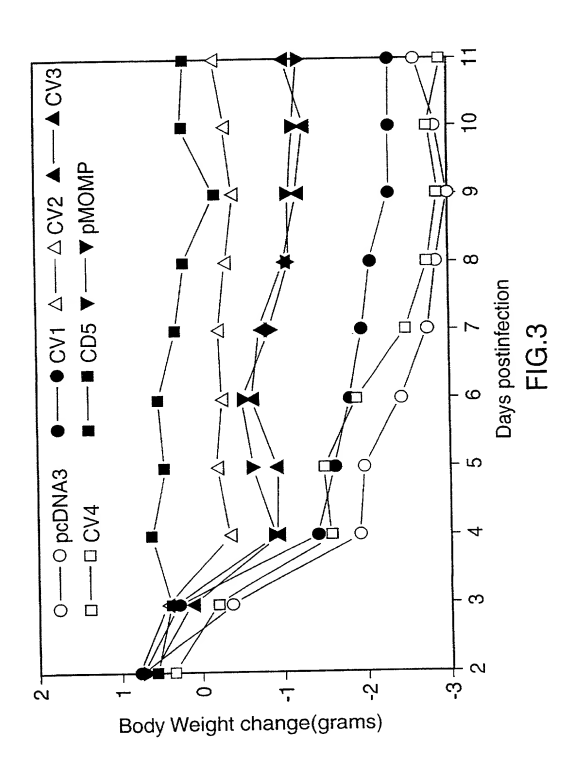
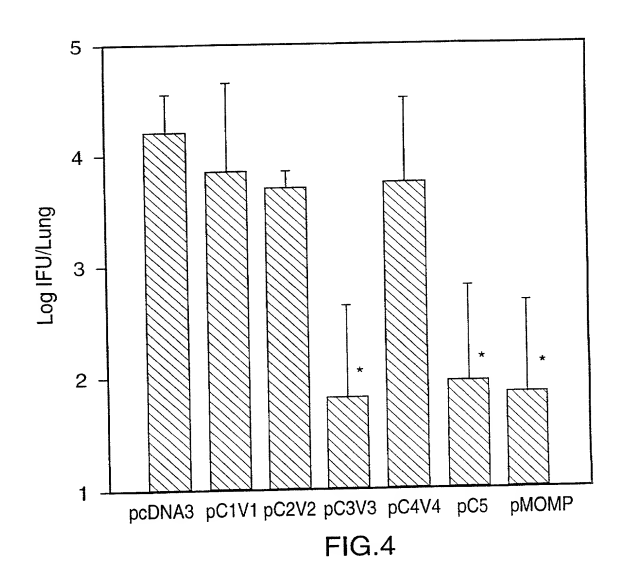
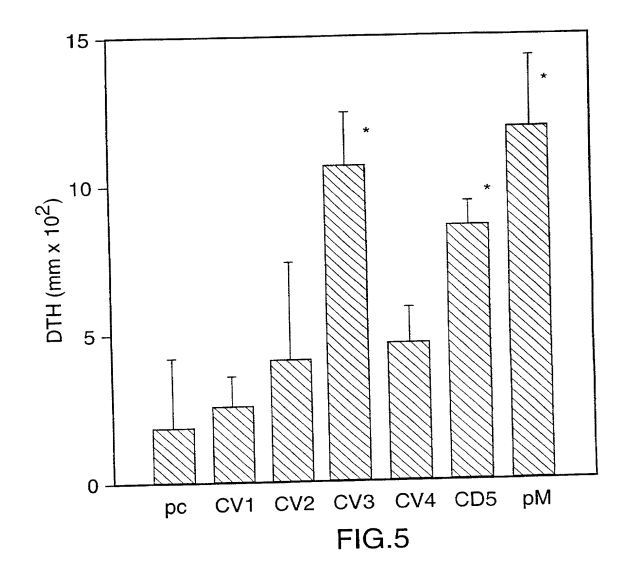


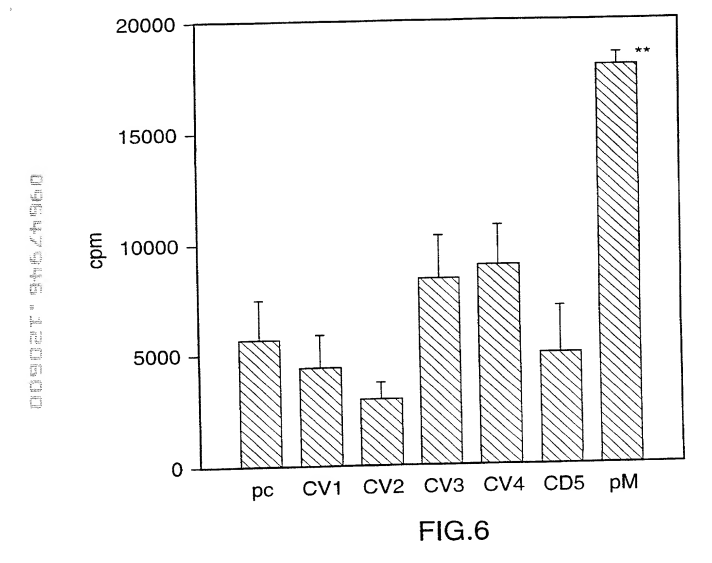
FIG.1



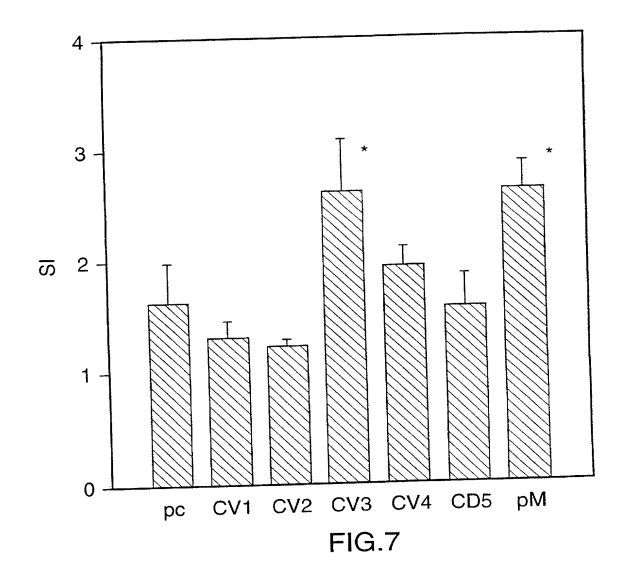


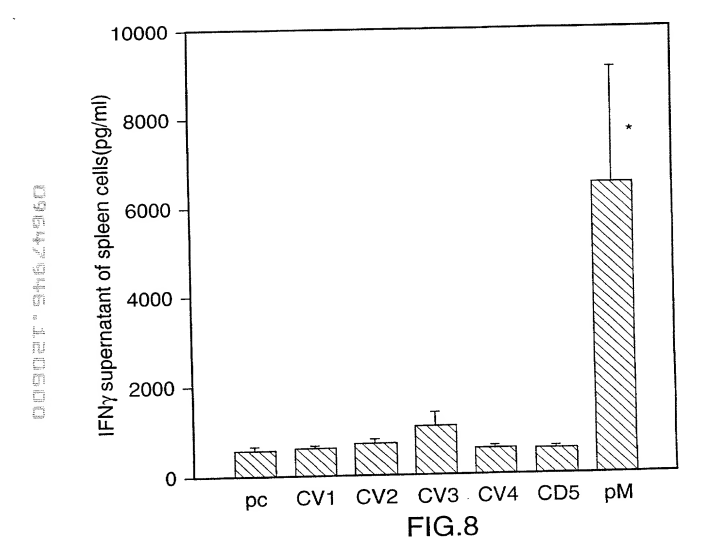




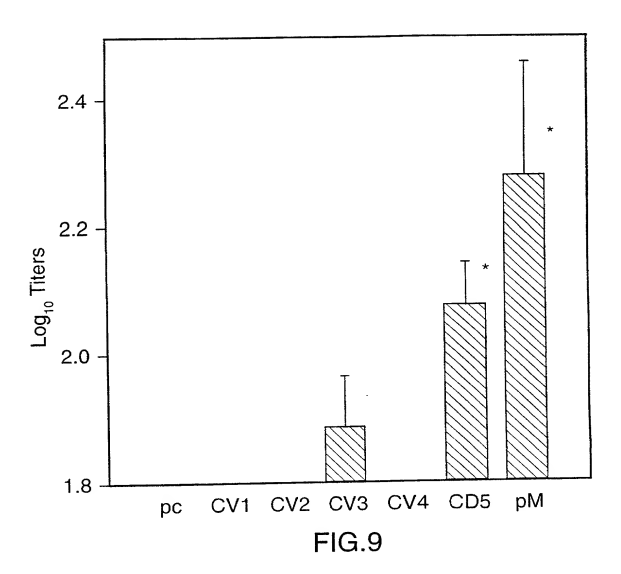




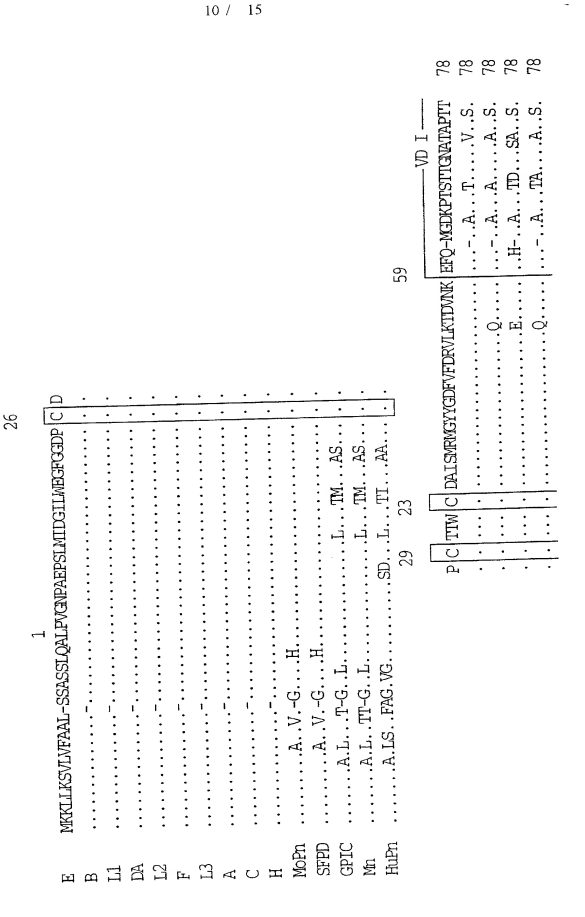




SUBSTITUTE SHEET (RULE 26)

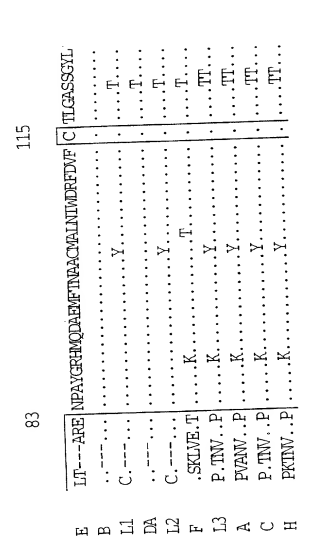






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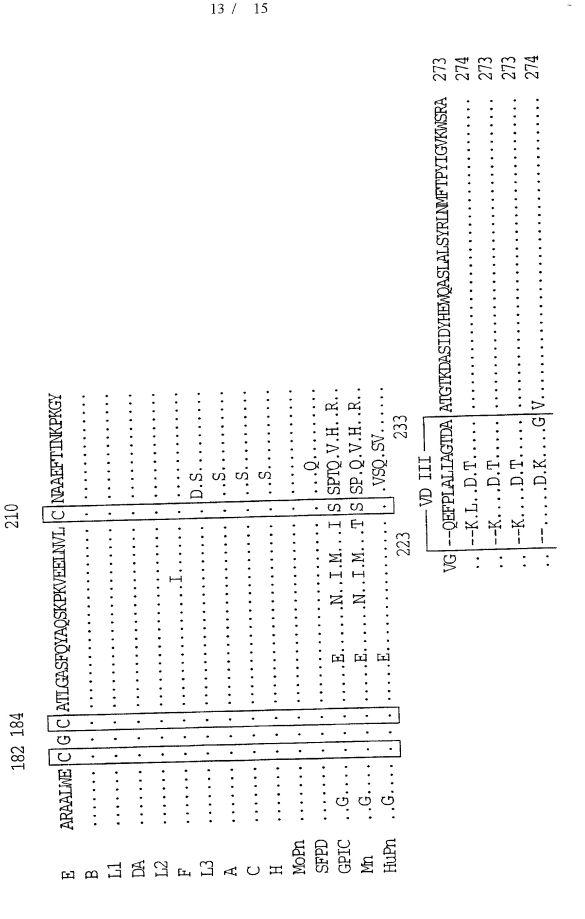


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FIG. 10D



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Declaration and Power of Attorney For Patent Application

English Language Declaration

As a below named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below next to

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled

the specification of which (check one) ☐ is attached hereto. ☒ was filed on April 7, 1999 as United States Application No. or PCT International Application Number PCT/CA99/00292							
☐ is attached hereto. ☑ was filed on April 7, 1999 as United States Application No. or PCT International Application Number PCT/CA99/00292							
was filed on April 7, 1999 as United States Application No. or PCT International Application Number PCT/CA99/00292							
Application Number PCT/CA99/00292							
and was amended on July 4, 2000							
(if applicable)							
I hereby state that I have reviewed and understand the contents of the above identified specification, including the claims, as amended by any amendment referred to above.							
I acknowledge the duty to disclose to the United States Patent and Trademark Office all information known to me to be material to patentability as defined in Title 37, Code of Federal Regulations, Section 1.56.							
I hereby claim foreign priority benefits under Title 35, United States Code, Section 119(a)-(d) or Section 365(b) of any foreign application(s) for patent or inventor's certificate, or Section 365(a) of any PCT International application which designated at least one country other than the United States, listed below and have also identified below, by checking the box, any foreign application for patent or inventor's certificate or PCT International application having a filing date before that of the application on which priority is claimed.							
Prior Foreign Application(s) Priority Not Claimed							
(Number) (Country) (Day/Month/Year Filed)							
(Number) (Country) (Day/Month/Year Filed)							
(Number) (Country) (Edy/Montas Four Floor)							
(Number) (Country) (Day/Month/Year Filed)							

application(s) listed below:	35 U.S.C. Section	119(e)	of any	United	States	provisional
	•					
(Application Serial No.)	(Filing Date)					
(Application Serial No.)	(Filing Date)					
(Application Serial No.)	(Filing Date)					

I hereby claim the benefit under 35 U. S. C. Section 120 of any United States application(s), or Section 365(c) of any PCT International application designating the United States, listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States or PCT International application in the manner provided by the first paragraph of 35 U.S.C. Section 112, I acknowledge the duty to disclose to the United States Patent and Trademark Office all information known to me to be material to patentability as defined in Title 37, C. F. R., Section 1.56 which became available between the filing date of the prior application and the national or PCT International filing date of this application:

09/055,765	April 7, 1998	Pending		
(Application Serial No.)	(Filing Date)	(Status) (patented, pending, abandoned)		
PCT/CA99/00292	April 7, 1999			
(Application Serial No.)	(Filing Date)	(Status) (patented, pending, abandoned)		
(Application Serial No.)	(Filing Date)	(Status) (patented, pending, abandoned)		

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

POWER OF ATTORNEY: As a named inventor, I hereby appoint the following attorney(s) and/or
agent(s) to prosecute this application and transact all business in the Patent and Trademark Office
connected therewith. (list name and registration number)
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_		

Second inventor's signature	Date
Residence	
Ditizenship	
Post Office Address	